

ds
>>>Unrecognizable Command
?ds
?ds
?ds

Set	Items	Description
S1	11905	EUKARYOTIC
S2	219809	EXPRESSION
S3	2094	S1 AND S2
S4	276006	BACTERIA OR COLI
S5	510	S3 AND S4
S6	7186	INTRON
S7	10	S5 AND S6 Euk. Exp. X Bact/coli intron
S8	0	SILENT (MUTATT (3W.) PROMOTER
S9	125	MUTATION (3W.) PROMOTER
S10	1	S8 AND S9 mut. pro
S11	703	LOW COPY
S12	3	S8 AND S11 low copy #

?ds s7
?s s7 or s10 or s12
10 S7
1 S10
3 S12
S13 14 S7 OR S10 OR S12

?d s13/3/1-14
>>>14.0 not recognized as item list

?d s13/3
<< Display 13/2/1
DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

11126778 BIOSIS Number: 97326778

Dihydrofolate reductase of Drosophila: Cloning and expression of a gene with a rare transcript

Hao H; Tyshenko M G; Walker V K

Dep. Biol., Queen's Univ., Kingston, ON K7L 3N6, CAN

Journal of Biological Chemistry 269 (21), 1994, 15175-15185.

Full Journal Title: Journal of Biological Chemistry

ISSN: 0021-9258

Language: ENGLISH

Print Number: Biological Abstracts Vol. 003 Iss. 003 Ref. 033200

Descriptors/Keywords: RESEARCH ARTICLE; DROSOPHILA; ESCHERICHIA COLI; EC 1.5.1.3; MOLECULAR SEQUENCE DATA; NUCLEOTIDE SEQUENCE; GENBANK-U06861; EMBL-U06861

Concept Codes:

*03500 Genetics and Cytogenetics-Animal

*10300 Replication, Transcription, Translation

--more--

?d s13/3/1-144
>>>Unrecognizable Command
?d s13/3/1-14

Display 13/3/1
DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

11126778 BIOSIS Number: 97326778

Dihydrofolate reductase of Drosophila: Cloning and expression of a gene with a rare transcript

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Hao H; Tyshenko M G; Walker V K
Dep. Biol., Queen's Univ., Kingston, ON K7L 3N6, CAN
Journal of Biological Chemistry 269 (21), 1994, 13173-13183.
Full Journal Title: Journal of Biological Chemistry
ISSN: 0021-9258
Language: ENGLISH

- end of record -

?
?d si3/3/2-14
(Display 13/3/2
DIALOG(R)File S:BIOSIS PREVIEW5(R)
(c) 1994 BIOSIS. All rts. reserv.

11038530 BIOSIS Number: 97238530
Genomic cloning of human thioredoxin-encoding gene: Mapping of the
transcription start point and analysis of the promoter
Kaghad M; Dessard F; Jacquemin-Gablon H; Caput D; Fradelizi D; Wollman E
E
Lab. INSERM U283, Pavillon Hardy A, Hopital Cochin, rue du Faubourg, St.
Jacques, 75671 Paris Cedex 14, FRA
Gene (Amsterdam) 140 (2), 1994, 273-278.
Full Journal Title: Gene (Amsterdam)
ISSN: 0378-1119
Language: ENGLISH

- end of record -

?d si3/3/3-14
>>>Unrecognizable Command
?t si3/3/3-144
>>>'1.4' not recognized as item list
?t si3/3/3-14

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13/3/3
DIALOG(R)File S:BIOSIS PREVIEW5(R)
(c) 1994 BIOSIS. All rts. reserv.

10830163 BIOSIS Number: 97030163
Cloning of the triosephosphate isomerase gene of Plasmodium falciparum
and expression in Escherichia coli
Ranie J; Kumar V P; Balaram H
Astra Res. Cent. India, P.O. Box 359, 10th Cross, Malleshwaram, Bangalore
560 003, IND
Molecular and Biochemical Parasitology 61 (2), 1993, 159-169.
Full Journal Title: Molecular and Biochemical Parasitology
ISSN: 0166-6851
Language: ENGLISH

13/3/4
DIALOG(R)File S:BIOSIS PREVIEW5(R)
(c) 1994 BIOSIS. All rts. reserv.

10428414 BIOSIS Number: 96028414
EUKARYOTIC TRANSLATION INITIATION FACTOR E FROM SACCCHAROMYCES-CEREVISIAE
CLONING CHARACTERIZATION AND EXPRESSION OF THE GENE ENCODING THE 43346 DA
PROTEIN
CHAKRAVARTI D; MAITRA
DEP. DEV. BIOL. CANCER, DIV. BIOL., ALBERT EINSTEIN COLL. MED., BRONX, NY
10461 USA

J BIOL CHEM 268 (14). 1993. 10524-10533. CODEN: JBCHIA
Full Journal Title: Journal of Biological Chemistry
Language: ENGLISH

13/3/5

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

10061557 BIOSIS Number: 95061557

EFFECTS OF INSERTIONS AND DELETIONS IN OLIGO NTRC OF ESCHERICHIA-COLI ON
NITROGEN REGULATOR I-DEPENDENT DNA BINDING AND TRANSCRIPTIONAL ACTIVATION
SHIAU S-P; CHEN P; REITZER L J
PROGRAM MOLECULAR CELL BIOL., UNIV. TEXAS DALLAS, P.O. BOX 830688,
RICHARDSON, TX 75083-0608, USA.

J BACTERIOL 175 (1). 1993. 190-199. CODEN: JOBAA
Full Journal Title: Journal of Bacteriology
Language: ENGLISH

13/3/6

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

9614625 BIOSIS Number: 94119625

GENETIC ANALYSIS OF THE HERPES SIMPLEX VIRUS TYPE 1 UL5 GENE ISOLATION OF
A LACZ INSERTION MUTANT AND EXPRESSION IN EUKARYOTIC CELLS
MALIK A K; MARTINEZ R; MUNCY L; CARMICHAEL C P; WELLER S K
DEP. MICROBIOL., UNIV. CONN. HEALTH CENT., FARMINGTON, CONN. 06030.
VIROLOGY 190 (2). 1992. 702-715. CODEN: VIRLA
Full Journal Title: Virology
Language: ENGLISH

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13/3/7

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

9569256 BIOSIS Number: 94074256

MOLECULAR CLONING STAGE-SPECIFIC EXPRESSION AND CELLULAR DISTRIBUTION OF
A PUTATIVE PROTEIN KINASE FROM PLASMODIUM FALCIPARUM
ZHAO Y; KAPPEL B; YANG J; FRANKLIN R M
DEP. STRUCTURAL BIOL., BIOCENT. UNIV. BASEL, KLINGELBERGSTRASSE 70,
CH-4056 BASEL, SWITZ.
EUR J BIOCHEM 207 (1). 1992. 305-313. CODEN: EJBDA
Full Journal Title: European Journal of Biochemistry
Language: ENGLISH

13/3/8

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

8105931 BIOSIS Number: 91026031

A MODULAR SET OF LAC-Z FUSION VECTORS FOR STUDYING GENE EXPRESSION IN
CAENORHABDITIS-ELIGANS
FIRE A; HARRISON S W; DIXON D
DEP. EMBRYOLOGY, CARNEGIE INST. WASHINGTON, 115 W. UNIVERSITY PKWY.,
BALTIMORE, MD. 21210, USA.

GENE (AMST) 93 (2). 1990. 135-150. CODEN: GENED

Full Journal Title: GENE (Amsterdam)
Language: ENGLISH

13/3/9

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

7647157 BIOSIS Number: 90015157

RNA POLYMERASE II TRANSCRIPTION BLOCKED BY ESCHERICHIA-COLI LAC REPRESSOR
DEUSCHLE U; HIPSCHIND R A; BUJARD H

ZENTRUM FUER MOLEKULARE BIOLOGIE, UNIVERSITÄT HEIDELBERG, INF 282.

D-6900 HEIDELBERG, W. GER.

SCIENCE (WASHINGTON D C) 248 (4954). 1990. 480-483. CODEN: SCICA

Full Journal Title: SCIENCE (Washington D C)

Language: ENGLISH

13/3/10

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

6602854 BIOSIS Number: 86069405

HOMOLOGOUS GENES FOR MOUSE 4.5S HYBRIDIZING RNA ARE FOUND IN ALL
EUKARYOTES AND THEIR LOW MOLECULAR WEIGHT RNA TRANSCRIPTS INTERMOLECULARLY
HYBRIDIZE WITH EUKARYOTIC 18S RIBOSOMAL RNA

TRINH-ROHLIK D; MAXWELL E S

DEP. BIOCHEMISTRY, BOX 7622, NORTH CAROLINA STATE UNIV., RALCIGH, NC
27695-7622.

NUCLEIC ACIDS RES 16 (13). 1988. 6041-6056. CODEN: NARHA

Full Journal Title: Nucleic Acids Research

Language: ENGLISH

(

13/3/11

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

5871681 BIOSIS Number: 84004246

ASSEMBLY OF THE MITOCHONDRIAL MEMBRANE SYSTEM MRPI AND MRP2 TWO YEAST
NUCLEAR GENES CODING FOR MITOCHONDRIAL RIBOSOMAL PROTEINS

MYERS A M; CRIVELLONE M D; TZAGOLOFF A

DEP. BIOCHEMISTRY AND BIOPHYSICS, IOWA STATE UNIV., AMES, IOWA 50011.

J BIOL CHEM 262 (7). 1987. 3380-3397. CODEN: JBCHA

Full Journal Title: Journal of Biological Chemistry

Language: ENGLISH

13/3/12

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

5447389 BIOSIS Number: 82092192

DNA-PROTEIN RECOGNITION DEMONSTRATION OF THREE GENETICALLY SEPARATED
OPERATOR ELEMENTS THAT ARE REQUIRED FOR REPRESSION OF THE ESCHERICHIA-COLI
DEO-CABD PROMOTERS BY THE DEO-R REPRESSOR

VALENTIN-HANSEN B A; LØRSEN J E L

DEP. MOLECULAR BIOL., ØRNSKJØLD UNIV., CAMPUSVEJ 55, 5230 ODENSE M, DEN.

EMBO (EUR MOL BIOL ORGAN) J 5 (6). 1986. 2015-2022. CODEN: EMJOD

Full Journal Title: EMBO (European Molecular Biology Organization)

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Journal
Language: ENGLISH

13/3/13

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

5275194 BIOSIS Number: 81042501

MOLECULAR CLONING AND REGULATED EXPRESSION OF THE HUMAN C-MYC GENE IN
ESCHERICHIA-COLI AND SACCHAROMYCES-CEREVISIAE COMPARISON OF THE PROTEIN
PRODUCTS

MIYAMOTO C; CHIZZONITE R; CROWL R; RUPPRECHT K; KRAMER R; SCHABER M;
KUMAR G; POONIAN M; JU G
DEP. MOLECULAR GENETICS, HOFFMANN-LA ROCHE, INC., ROCHE RES. CENT.,
NUTLEY, N.J. 07110.

PROC NATL ACAD SCI U S A 82 (21). 1985. 7232-7236. CODEN: PNASA

Full Journal Title: Proceedings of the National Academy of Sciences of
the United States of America

Language: ENGLISH

13/3/14

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

4024878 BIOSIS Number: 75072237

BOTH EARLY AND LATE CONTROL SEQUENCES OF SV-40 AND POLYOMA PROMOTE
TRANSCRIPTION OF ESCHERICHIA-COLI GPT GENE IN TRANSFECTED CELLS

BOURACHOT B; JOUANNEAU J; GIRI I; KATINKA M; CERESINI S; YANIV M
UNITE VIRUS ONCOGENES, DEP. BIOL. MOL., INST. PASTEUR, 25 RUE DR. ROUX,
75724 PARIS CEDEX 15, FR.

EMBO (EUR MOL BIOL ORGAN) J 1 (8). 1982. 895-900. CODEN: EMJOD

Full Journal Title: EMBO (European Molecular Biology Organization)
Journal

Language: ENGLISH

?ds{>>>Unrecognizable Command

?t si4/3/i-12

>>>Unrecognizable Command

?{t{ s

>>>Unrecognizable Command

?t si4/3/i-12

{

14/3/1

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

11295317 BIOSIS Number: 07405317

Expression and export in Escherichia coli of fusion proteins containing
carboxy-terminally located honeybee preproinsulin

He M; Liu H; Austen B

Dep. Surgerv, St. George's Hosp. Med. Sch., Cranmer Terrace, London SW17
ORE, UK

DNA and Cell Biology 13 (8). 1994. 875-882.

Full Journal Title: DNA and Cell Biology

ISSN: 1044-5408

Language: ENGLISH

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14/3/2

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

11221285 BIOSIS Number: 97421285

A first step in the development of gene therapy for colorectal carcinoma:
Cloning, sequencing, and expression of Escherichia coli cytosine deaminase
Austin E A; Huber D E
Div. Cell Biology, Wellcome Res. Labs., 3030 Cornwallis Road, Research
Triangle Park, NC 27709, USA
Molecular Pharmacology 43 (3), 1993, 380-387.
Full Journal Title: Molecular Pharmacology
ISSN: 0026-895X
Language: ENGLISH

14/3/3

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

11203145 BIOSIS Number: 97403145

Escherichia coli ppt as a positive selectable marker in embryonal stem
cells
Spring K J; Mattick J S; Don R H
Centre Mol. Biol. Biotechnol., Univ. Queensland, Brisbane, QLD 4072, AUL
Biochimica et Biophysica Acta 1218 (2), 1994, 155-162.
Full Journal Title: Biochimica et Biophysica Acta
ISSN: 0006-3002
Language: ENGLISH

14/3/4

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

10898047 BIOSISNumber: 97098047

Protection against chloroethylnitrosourea cytotoxicity by eukaryotic
3-methyladenine DNA glycosylase
Matijasevic Z; Doosalis M; Mackay W; Samson L; Ludlum D B
Dep. Pharmacol., Univ. Massachusetts Med. Sch., Worcester, MA 01655, USA
Proceedings of the National Academy of Sciences of the United States of
America 90 (24), 1993, 11855-11859.
Full Journal Title: Proceedings of the National Academy of Sciences of
the United States of America
ISSN: 0027-0424
Language: ENGLISH

14/3/5

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

9142352 BIOSIS Number: 93127352

OVEREXPRESSION OF HIGHER EUKARYOTIC MEMBRANE PROTEINS IN BACTERIA NOVEL
INSIGHTS OBTAINED WITH THE LIVER MITOCHONDRIAL PROTON PHOSPHATE SYMPORTER
FERREIRA G C; PEDERSEN P L
LAB. MOL. CELLULAR BIOENERGETICS, DEP. BIOL. CHEM., JOHNS HOPKINS UNIV.,
SCH. MED., BALTIMORE, MD 21205.
J BIOL CHEM 267 (8), 1992, 5400-5460. CODEN: JBIOA
Full Journal Title: Journal of Biological Chemistry
Language: ENGLISH

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14/3/6

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

(c) 1994 BIOSIS. All rts. reserv.

9138161 BIOSIS Number: 93123161

DICTYOSTELIUM-DISCOIDEUM AS AN EXPRESSION HOST FOR THE CIRCUMSPOROZOITE
PROTEIN OF PLASMODIUM-FALCIPARUM

FASEL N; BEGDADI-RAIS C; BERNARD M; BRON C; CORRADIN G; REYMOND C D

ISREC, CH-1066 EPALINGES, SWITZ.

GENE (AMST) 111 (2), 1992, 157-163. CODEN: GENED

Full Journal Title: GENE (Amsterdam)

Language: ENGLISH

14/3/7

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

(c) 1994 BIOSIS. All rts. reserv.

9094147 BIOSIS Number: 93079147

DIPHThERIA TOXIN RECEPTOR-BINDING DOMAIN SUBSTITUTION WITH INTERLEUKIN &
GENETIC CONSTRUCTION AND INTERLEUKIN 6 RECEPTOR-SPECIFIC ACTION OF A
DIPHThERIA TOXIN-RELATED INTERLEUKIN 6 FUSION PROTEIN

JEAN L-F L; MURPHY J R

EVANS DEP. CLIN. RES., DEP. MED., UNIV. HOSP., 88 EAST NEWTON ST.,
BOSTON, MASS. 02118.

PROTEIN ENG 4 (8), 1991, 989-994. CODEN: PRENC

Full Journal Title: Protein Engineering

Language: ENGLISH

14/3/8

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

(c) 1994 BIOSIS. All rts. reserv.

9079316 BIOSIS Number: 93064316

TRANSFER OF THE BACTERIAL GENE FOR CYTOSINE DEAMINASE TO MAMMALIAN CELLS
CONFERS LETHAL SENSITIVITY TO 5 FLUOROCYTOSINE A NEGATIVE SELECTION SYSTEM

MULLEN C A; KILSTRUP M; BLAESE R M

CELLULAR IMMUNOLOGY SECTION, METABOLISM BRANCH, NATL. CANCER INST., NATL.
INST. HEALTH, BUILDING 10, ROOM 6B05, BETHESDA, MD. 20892.

PROC NATL ACAD SCI U S A 89 (1), 1992, 33-37. CODEN: PNASA

Full Journal Title: Proceedings of the National Academy of Sciences of
the United States of America

Language: ENGLISH

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14/3/9

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

(c) 1994 BIOSIS. All rts. reserv.

8627136 BIOSIS Number: 92092136

IN-VITRO ACTIVATION OF ESCHERICHIA-COLI PROMENOLYSIN TO THE MATURE
MEMBRANE-TARGETED TOXIN REQUIRES HLY-C AND A LOW MOLECULAR-WEIGHT CYTOSOLIC
POLYPEPTIDE

HARDIE K R; ISCARTEL J P; KORONAKIS E; HUGHES E; KORONAKIS V

DEP. PATHOL., UNIV. CAMB., TENNIS COURT ROAD, CAMBRIDGE CB2 1QP, UK.

MOL MICROBIOL 5 (7), 1991, 1665-1680. CODEN: MONTE

Full Journal Title: Molecular Microbiology

Language: ENGLISH

14/3/10

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

7749587 BIOSIS Number: 90117587

ON THE MODE OF ACTION OF BASIC PHOSPHOLIPASE A-2 FROM NAJA-NIGRICOLLIS
VENOM

CHWETZOFF S

CET. D'ETUDES NUCL. DE SACLAY, SERV. BIOCHIM., BAT. 142. 91191
GIF-SUR-YVETTE CEDEX, FR.

BIOCHIM BIOPHYS ACTA 1045 (3). 1990. 285-290. CODEN: BBACA

Full Journal Title: Biochimica et Biophysica Acta

Language: ENGLISH

14/3/11

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

7635725 BIOSIS Number: 90003725

PHENOTYPIC SELECTION AND CHARACTERIZATION OF MUTANT ALLELES OF A
EUKARYOTIC DNA TOPOISOMERASE I

MORHAM S G; SHUMAN S

PROGRAM MOLECULAR BIOLOGY, SLOAN-KETTERING INST., NEW YORK, N.Y. 10021.

GENES DEV 4 (4). 1990. 515-524. CODEN: GEDEE

Full Journal Title: Genes & Development

Language: ENGLISH

14/3/12

DIALOG(R)File 5:BIOSIS PREVIEWS(R)
(c) 1994 BIOSIS. All rts. reserv.

6613314 BIOSIS Number: 86079865

THE INSTABILITY OF A RECOMBINANT PLASMID CAUSED BY A PROKARYOTIC-LIKE
PROMOTER WITHIN THE EUKARYOTIC INSERT CAN BE ALLEVIATED BY EXPRESSION OF
ANTISENSE RNA

FUTTERER J; GORDON K; PFEIFFER P; HOHN T

FRIEDRICH MIESCHER INSTITUT, P.O. BOX 2543, CH-4002 BASEL, SWITZERLAND.

GENE (AMST) 67 (1). 1988. 141-145. CODEN: GENED

Full Journal Title: GENE (Amsterdam)

Language: ENGLISH

?{({{

>>>Unrecognizable Command

?ds

Set	Items	Description
51	11905	EUKARYOTIC
52	219809	EXPRESSION
53	2094	51 AND 52
54	276006	BACTERIA OR COLI
55	516	53 AND 54
56	7186	INTRON
57	10	55 AND 56
58	0	SILENT () TAT? (3W.) PROMOTER
59	125	MUTATION (3W.) PROMOTER
510		55 AND 56

BEST AVAILABLE COPY

S11 763 LOW(COPY)
 S12 3 55 AND S11
 S13 14 57 OR S13 OR S12
 S14 12 55 AND TOXIC?
 ?(d d
 {?d s10/3/i
 >>>Unrecognizable Command
 ?d{
 >>>Unrecognizable Command
 ?d s10/12/i
 >>>'12' not a valid format name
 ?d s110
 { Display 14/2/i
 DIALOG(R)File S:BIOSIS PREVIEW(R)
 (c) 1994 BIOSIS. All rts. reserv.

11295317 BIOSIS Number: 97495317
 Expression and export in Escherichia coli of fusion proteins containing
 carboxy-terminally located honeybee preproinsulin
 He M; Liu H; Austen B
 Dep. Surgery, St. George's Hosp. Med. Sch., Cranmer Terrace, London SW17
 ORE. UK
 DNA and Cell Biology 13 (3). 1994. 875-882.
 Full Journal Title: DNA and Cell Biology
 ISSN: 1044-5498
 Language: ENGLISH
 Print Number: Biological Abstracts Vol. 090 Iss. 010 Ref. 131056
 Descriptors/Keywords: RESEARCH ARTICLE; ESCHERICHIA COLI
 Concept Codes:
 *03300 Genetics and Cytogenetics-Animal
 *10062 Biochemical Studies Nucleic Acids, Purines and Pyrimidines
 *10064 Biochemical Studies-Proteins, Peptides and Amino Acids

- more -

{?d s10/3/i
 >>>Unrecognizable Command
 ?d s10/3/i
 { Display 10/3/i
 DIALOG(R)File S:BIOSIS PREVIEW(R)
 (c) 1994 BIOSIS. All rts. reserv.

10061557 BIOSIS Number: 95061557
 EFFECTS OF INSERTIONS AND DELETIONS IN GLNG NTRC OF ESCHERICHIA-COLI ON
 NITROGEN REGULATOR I-DEPENDENT DNA BINDING AND TRANSCRIPTIONAL ACTIVATION
 SHIAU S-P; CHEN P; REITZER L J
 PROGRAM MOLECULAR CELL BIOL., UNIV. TEXAS DALLAS, P.O. BOX 830688,
 RICHARDSON, TX 75083-0688, USA.
 J BACTERIOL 175 (1). 1993. 190-199. CODEN: JODAA
 Full Journal Title: Journal of Bacteriology
 Language: ENGLISH

- end of display -

?s review
 S15 240332 REVIEW
 ?{
 >>>Unrecognizable Command
 ?s s4 and s2
 { 270000 50
 210000 50
 S1A 10001 12 AND 17

BEST AVAILABLE COPY

```

?s si6 and si5
>>>Unrecognizable Command
?(s si6 and si5
>>>Unrecognizable Command
>>>Excessive invalid command rate. Session Terminated.
25nov94 11:29:57 User217744 Session D351.2
    $16.96    @.316 Hrs File3
        $2.00  2 Type(s) in Format  2
        $27.00 27 Type(s) in Format  3
    $29.00  29 Types
        $7.00  View Fee
    $54.96 Estimated cost File3
    $3.60  TYMNET
    $58.56 Estimated cost this search
    $58.09 Estimated total session cost    @.328 Hrs.
Logoff: level 37.11.01 D 11:29:57

```

TYMNET: call cleared by request

please log in: dialog

DIALOG: call connected

DIALOG INFORMATION SERVICES

PLEASE LOGON:

?0000000000

ENTER PASSWORD:

?0000000000

Welcome to DIALOG

Dialog level 37.11.01D

BEST AVAILABLE COPY

Last logoff: 25nov94 11:29:57

Logon file405 25nov94 11:30:20

SYSTEM:HOME

Menu System II: D2 version 1.7.1 term=ASCII

*** DIALOG HOMEDBASE(SM) Main Menu ***

Information:

1. Announcements (new files, free connect time, price changes, etc.)
2. Database, Rates, & Command Descriptions
3. Help in Choosing Databases for Your Topic
4. Customer Services (telephone assistance, training, seminars, etc.)
5. Product Descriptions

Connections:

6. DIALOG Menus(SM)
7. DIALOG Business Connection(R), Headlines(SM), Medical Connection(SM)
8. DIALOG SourceOne(SM) Document Delivery
9. Data-Star
10. Other Online Menu Services & Files (MoneyCenter(R), DAG, TNT, etc.)

/H = Help

/L = Logoff

/NOMENU = Command Mode

Enter an option number to view information or to connect to an online service. Enter a DECIN command plus a file number search a database (e.g., B1 for ERIC).

265

(25nov94 11:30:45 User217744 Session D352.1
 \$0.00 0.007 Hrs FileHomeBase
 \$0.00 Estimated cost FileHomeBase
 \$0.00 TYMNET
 \$0.00 Estimated cost this search
 \$0.00 Estimated total session cost 0.007 hrs.

File 5:BIOSIS PREVIEWS(R) 1969-1994/Dec W3
 (c) 1994 BIOSIS

Set Items Description

?s {pro
 >>>Unrecognizable Command
 ?s{ prokaryotic()expression
 0 .(PROKARYOTIC
 219809 EXPRESSION
 S1 0 .(PROKARYOTIC()EXPRESSION

?s proka(rv
 >>>Null command ignored
 ?s prokaryotic()expression
 3152 PROKARYOTIC
 219809 EXPRESSION
 S2 131 PROKARYOTIC()EXPRESSION

?s review
 S3 240332 REVIEW

?s s2 and s3
 131 S2
 240332 S3
 S4 2 S2 AND S3-

?s coli(3w)expression
 156325 COLI
 0 E.XPRESSION
 S5 0 COLI(3W)E.XPRESSION

?s coli(3w)expression
 156325 COLI
 219809 EXPRESSION
 S6 1665 COLI(3W)EXPRESSION

?s s6 and s3
 1665 S6
 240332 S3
 S7 13 S6 AND S3

?
 >>>Unrecognizable Command
 ?s s4 or s7

2 S4
 13 S7
 S8 15 S4 OR S7

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DIALOG(R)File 5:BIOSIS PREVIEWS(R)
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11312250 BIOSIS Number: 07512250

Cloning of a wheat 13-kDa grain softness protein (GSP) GSP is a mixture of purindoline-like polypeptides

Rahman S; Jolly C J; Chen H; J H; Hallowell A

Canadian Health Scientific Institute, 300, University Ave, Saint

Prok wpx Review on Coli Exp. & Review

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Industry, P.O. Box 1600, Australian Capital Territory 2601, AUL

European Journal of Biochemistry 223 (3), 1994, 917-925.

Full Journal Title: European Journal of Biochemistry

ISSN: 0014-2956

Language: ENGLISH

Print Number: Biological Abstracts Vol. 298 Iss. 011 Ref. 147989

Descriptors/Keywords: RESEARCH ARTICLE; STARCH PROTEIN; LIPID BINDING
PROPERTIES; GRAIN SOFTNESS; CULTIVAR VARIATION; GRAIN QUALITY; MOLECULAR
SEQUENCE DATA; NUCLEOTIDE SEQUENCE; AMINO ACID SEQUENCE

Concept Codes:

- *03504 Genetics and Cytogenetics-Plant
- *10064 Biochemical Studies-Proteins, Peptides and Amino Acids
- *10506 Biophysics-Molecular Properties and Macromolecules
- *13210 Nutrition-Water-Soluble Vitamins
- *13530 Food Technology-Evaluations of Physical and Chemical Properties
(1970-)
- *51512 Plant Physiology, Biochemistry and Biophysics-Reproduction
- *51522 Plant Physiology, Biochemistry and Biophysics-Chemical
Constituents
- *52504 Agronomy-Grain Crops
- 10066 Biochemical Studies-Lipids
- 10068 Biochemical Studies-Carbohydrates

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Biosystematic Codes:

25305 Gramineae

Super Taxa:

Plants; Vascular Plants; Spermatophytes; Angiosperms; Monocots

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DIALOG(R)File 5:BIOSIS PREVIEW(R)

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11312250 BIOSIS Number: 97512250

Cloning of a wheat 15-kDa grain softness protein (GSP) GSP is a mixture
of puroindoline-like polypeptides

Rahman S; Jolly C J; Skerritt J H; Walloscheck A

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European Journal of Biochemistry 223 (3), 1994, 917-925.

Full Journal Title: European Journal of Biochemistry

ISSN: 0014-2956

Language: ENGLISH

The wheat starch 15-kDa protein (called grain softness protein or GSP) consists of a major polypeptide and several minor polypeptides. An antiserum raised against GSP was used to screen a wheat cDNA library. A cDNA family encoding approximately 15-kDa proteins that included a heptapeptide sequence previously isolated from protease digests of GSP was identified. A partial cDNA was used in a prokaryotic expression system to produce a fusion protein which reacted strongly against the original anti-GSP serum. A new antiserum raised against the fusion protein produced a weak reaction against a 15-kDa polypeptide extracted from wheat seeds. The results suggest that the proteins encoded by the cDNA family form a minor component of the mixture of 15-kDa polypeptides defined as GSP. RNA complementary to the cDNAs could be extracted from both soft and hard wheat grains from about half-way through grain filling. The encoded proteins are novel members of the 2S superfamily of seed proteins, a diverse family of proteins which maintain a characteristic framework of cysteine residues. The deduced proteins show the highest similarity to the oat 16-kDa avenin and to wheat puroindoline (a lipid-binding 15-kDa protein from wheat). Review of previously published data shows that puroindoline is also closely

related to the major polypeptide of GSP, suggesting that the lipid-binding properties of GSP polypeptides may influence brain softness.

8/7/2

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11271932 BIOSIS Number: 97471932

Pattern and control in bacterial colony development

Shapiro J A

Dep. Biochem. and Molecular Biol., Univ. Chicago, 920 E. 58th St.,
Chicago, IL 60637, USA

Science Progress 76 (301-302), 1992, 399-424.

Full Journal Title: Science Progress

ISSN: 0036-8504

Language: ENGLISH

8/7/3

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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11248095 BIOSIS Number: 97448095

The molecular biology of production cell lines

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Biologicals 22 (2), 1994, 85-94.

Full Journal Title: Biologicals

ISSN: 1045-1056

Language: ENGLISH

The emergence of a wide variety of biological expression systems for the large-scale production of therapeutic proteins has shifted the focus from vectors to host organisms. Although expression systems now span bacteria, fungi, plants, insects, and mammalian cells, the vast majority of recombinant-derived biopharmaceuticals at the present time have been produced in *Escherichia coli* and in mammalian cells. This promises to change as the economic benefits of the newer systems permit the development of a new generation of proteins heretofore considered unfeasible for commercial development. Despite the impressive results which have been observed for many of the newer systems, there are many commercial considerations which suggest that *E. coli* and CHO cell expression systems may continue to dominate the manufacture of biopharmaceuticals for a long time to come.

8/7/4

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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11029432 BIOSIS Number: 97229432

The cold-shock response-a hot topic

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Molecular Microbiology 11 (5), 1994, 811-818.

Full Journal Title: Molecular Microbiology

ISSN: 0950-382X

Language: ENGLISH

The cold-shock response of *Escherichia coli* involves a specific pattern

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of gene expression in response to abrupt shifts to lower temperatures. This pattern includes the induction of cold-shock proteins, synthesis of proteins involved in transcription and translation, and repression of heat-shock proteins. The identified cold-shock proteins are involved in various cellular functions from supercoiling of DNA to initiation of translation. The major cold-shock protein, CspA, has high sequence similarity with three other *E. coli* proteins - CspB, CspC, and CspD. Using translational lacZ fusions, cspB was found to be cold-shock inducible at the level of transcription like cspA, while cspC and cspD were not. The Csp proteins, which share sequence similarity with other prokaryotic proteins and with the 'coldshock domain' of eukaryotic Y-box proteins, may have a function in activating transcription or unwinding or masking RNA molecules. Because the coldshock response can also be induced by the addition of certain inhibitors of translation, it has been proposed that the state of the ribosome is the physiological sensor for the induction. In addition to *E. coli*, cold-shock proteins have also been found in other prokaryotic and eukaryotic organisms.

8/7/5

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10804882 BIOSIS Number: 07004882

Expression of modified cytochrome P450 2C10 (2C7) in *Escherichia coli*. purification, and reconstitution of catalytic activity

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Nashville, TN 37232-0146, USA

Archives of Biochemistry and Biophysics 300 (2), 1993. 443-450.

Full Journal Title: Archives of Biochemistry and Biophysics

ISSN: 0003-9861

Language: ENGLISH

The human cytochrome P450 (P450) 2C gene family is complex and heterologous expression methods are needed to facilitate the isolation of individual P450 proteins and the elucidation of their catalytic specificities. We prepared a series of constructs of P450 2C10 in the plasmid vector pCW, with modification of the 5' end of the coding sequence of the cDNA. Some were not expressed at all in *Escherichia coli*; two were expressed at levels of 5-20 nmol membrane-bound P450 (liter culture)-i-one (2C1028) with original codons 2-7 altered by substitution of the 5'-terminal sequence described by Barnes et al. (Barnes, H. J., Ariotto, M. P., and Waterman, M. R., Proc. Natl. Acad. Sci. USA 88, 5597-5601, 1991) and one (2C1029) with original codon 2 modified, codons 3-20 deleted, and alteration of the immediate downstream codons. In both cases the P450 2C10 proteins were found essentially only in the bacterial membranes. These proteins could be purified to a high degree by solubilization and a single DEAE chromatography step. Typical P450 Fe-2+ centdot CD absorption spectra were observed in the bacterial membranes and the purified preparations. The P450 2C1029 protein was found to have its N-terminal Met removed and the expected residues 2 (Ala)-24 were identified by amino acid sequence analysis. However, the other P450 (2C1028) was apparently blocked at the N-terminus. Three native P450 2C7/10 preparations isolated from human liver showed the expected sequences (beginning with Met) for at least the first 17 residues. The blocked N-terminus in the P450 2C1028 protein may be the result of the MALLAVT sequence, which was also used in the expression of P450 3A4 and resulted in a blocked protein. Catalytic activities of P450 2C1028 and P450 2C1029 to tolbutamide hydroxylation were similar to those measured with purified liver P450 2C7/10 in the presence of cytochrome b 5. Although the effect of cytochrome b 5 did not always affect the same pattern

as with the isolated liver enzyme. The recombinant P450 2C10 enzymes did not catalyze (S)-mephenytoin 4'-hydroxylation.

8/7/6

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10005590 BIOSIS Number: 95005590

CAPSULES OF ESCHERICHIA-COLI EXPRESSION AND BIOLOGICAL SIGNIFICANCE

JANN K; JANN B

MAX-PLANCK-INSTITUT IMMUNBIOLOGIE, STUEBEWEG 51, FREIBURG-ZACHRINGEN, GERMANY.

CAN J MICROBIOL 38 (7). 1992. 705-710. CODEN: CJMIA

Full Journal Title: Canadian Journal of Microbiology

Language: ENGLISH

Escherichia coli may cause intestinal or extraintestinal infections. Generally, extraintestinal E. coli are encapsulated. The capsules are important virulence determinants, which enable the pathogenic bacteria to evade or counteract the unspecific host defense during the early (preimmune) phase of infection. They interfere with the action of complement and phagocytes. This effect is generally transient and overcome by capsule-specific antibodies in the immune phase of the host defense. In some cases, capsules are not or only poorly immunogenic, as a result of structural relationship or identify with host material. Strains with such capsules (e.g., K1 or K5) are very virulent. Bacterial capsules consist of acidic polysaccharides, which are made up from oligosaccharide repeating units. The capsules of E. coli are divided into two groups, which differ in chemistry, biochemistry, and genetic organization. All capsular polysaccharides are chromosomally determined: those of group I close to his and those of group II close to serA. The biosynthesis and surface expression have been extensively studied with representatives of group II capsular polysaccharides. It could be shown that their biosynthesis is directed from a pene block that determines the synthesis of the polysaccharide, its translocation across the cytoplasmic membrane, as well as its surface expression in a coordinate process. The chemical nature of group II capsular polysaccharides, as well as the mechanism(s) of their biosynthesis and expression, is presented.

8/7/7

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9795891 BIOSIS Number: 44045891

BACTERIOPHAGE LAMBDA AS A CLONING VECTOR

CHAUTHAIWALE V M; THERWATH A; DESHPANDE V V

DIV. BIOCHEM. SCI., NATIONAL CHEM. LAB., PUNE 411 005, INDIA.

MICROBIOL REV 56 (4). 1992. 577-591. CODEN: MBRED

Full Journal Title: Microbiological Reviews

Language: ENGLISH

8/7/8

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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9560655 BIOSIS Number: 94006555

RECOMBINANT GRANULOCYTE-MACROPHAGE COLONY-STIMULATING FACTOR FROM CSF A

STUDY OF ITS PHARMACOLOGICAL PROPERTIES AND EFFECTIVE ROLE IN THE

MANAGEMENT OF MYELOSUPPRESSION

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ADIS INTERNATIONAL LIMITED, 41 CENTORIAN DRIVE, PRIVATE BAG 65061,
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DRUGS 43 (4), 1992, 510-560. CODEN: DRUGA

Full Journal Title: Drugs

Language: ENGLISH

Recombinant granulocyte-macrophage colony stimulating factor (rGM-CSF) is a polypeptide hormone produced through recombinant DNA technologies in glycosylated (yeast or mammalian expression systems) or nonglycosylated (*Escherichia coli* expression system) form. It is a multilineage haematopoietin which stimulates proliferation and differentiation of bone marrow myeloid progenitors and increases peripheral white blood cell counts when administered systemically. Treatment is generally well tolerated, although mild to moderate flu-like symptoms are common and rGM-CSF-induced fever and fluid retention may be problematic in occasional patients. rGM-CSF accelerates recovery of peripheral neutrophil counts after bone marrow transplantation, and results of a placebo-controlled randomised trial correlate this with reduced infectious episodes and shortened length of hospitalisation in patients with lymphoid malignancies. A substantial number of patients with graft failure after bone marrow transplantation also respond to rGM-CSF. The duration of myelosuppression secondary to cancer chemotherapy can be significantly reduced by rGM-CSF which has permitted investigation of antineoplastic dose-intensity escalation. In some haematopoietic disorders (e.g. aplastic anaemia, myelodysplasia and neutropenia secondary to HIV infection and antiviral therapy), rGM-CSF produces clinically useful increases in peripheral blood granulocyte counts, although the effect is generally not sustained after drug withdrawal. The potential for rGM-CSF to stimulate proliferation of the abnormal clone in myelodysplasia and in acute myelogenous leukaemia following induction therapy is of concern. Available data suggest, however, that with appropriate monitoring and exclusion of high-risk patients this serious potential risk can be avoided, and that myelopoiesis is enhanced in such patients by rGM-CSF treatment. Recombinant colony-stimulating factors are a new therapeutic modality; hence many aspects of their use remain to be clarified. Nonetheless, as one of a small group of novel agents rGM-CSF has major potential in the management of myelosuppression secondary to cytoreductive therapy with or without bone marrow transplantation, and in amelioration of disturbed myelopoiesis. It represents an important application of biotechnology to a difficult area of therapeutics.

8/7/9

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9512370 BIOSIS Number: 94017370

ANTIBODY ENGINEERING THE USE OF *ESCHERICHIA-COLI* AS AN EXPRESSION HOST
WARD E S

CANCER IMMUNOBIOLOG. CENT., UNIV. TEXAS SOUTHWESTERN MED. CENTER, DALLAS,
TEX. 75235-9048, USA.

FASEB (FED AM SOC EXP BIOL) J 6 (7), 1992, 2422-2427. CODEN: FAJDE

Full Journal Title: FASEB (Federation of American Societies for
Experimental Biology) Journal

Language: ENGLISH

The hypervariable loops of an antibody molecule are supported on the relatively conserved beta-sheeted framework of the heavy and light-chain variable domains (designated VH and VL domains, respectively). Residues within and flanking these loops interact with antigen and confer the specificity and affinity of antigen binding on the immunoglobulin

molecule. Thus, the isolation and expression of VH and VL domain genes are of particular interest both for analysis of the determinants of antibody specificity and for generation of fragments with binding affinities for use in therapy and diagnosis. The PCR can now be used to isolate diverse repertoires of antibody VH and VL domain genes from antibody-producing cells from different species, including humans and mice. The genes can be expressed as either secreted or surface-bound Fv or Fab fragments, using Escherichia coli expression systems, and the desired antigen-binding specificity screened for or, preferably, selected. The use of E. coli as an expression host allows the required antigen-binding specificity to be isolated in clonal form in a matter of days. The VH and VL domain genes can also be hypermutated and higher-affinity variants isolated by screening or selection. Thus, the use of this technology should allow the isolation of novel binding specificities or specificities that are difficult to generate by hybridoma technology. It will also facilitate the isolation of human-derived Fv/Fab fragments that may be less immunogenic in therapy. This approach therefore has almost unlimited potential in the generation of therapeutics with binding of specificities to order. The fragments can be used either alone or linked to effector functions in the form of antibody-constant domains or toxins. The new technology could prove to be a method of choice for rapid and convenient production of designer antibodies.

8/7/10

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8196807 BIOSIS Number: 91117807

PROTEIN OVERPRODUCTION FOR ORGANIC CHEMISTS

SCHREIBER S L; VERDINE G L

DEP. CHEM., HARVARD UNIV., CAMBRIDGE, MASS. 02138.

TETRAHEDRON 47 (14-15), 1991, 2543-2562. CODEN: TETRA

Full Journal Title: Tetrahedron

Language: ENGLISH

In this review we present the principles behind protein overexpression in bacteria, emphasizing how this biosynthetic system can be manipulated to generate large quantities of proteins for study. In addition to the classical (molecular biological), methods for constructing protein-overproducing bacterial, we discuss our recently developed (chemical/enzymatic) method, the Expression Cassette polymerase Chain Reaction (ECPCR). The chemical/enzymatic transformation of an unexpressable to an expressable gene afforded by ECPCR can routinely be carried out in the experimental organic chemistry laboratory, hence, ECPCR offers a convenient point of entry for chemists interested in macromolecular science.

((

8/7/11

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7842550 BIOSIS Number: 40043550

FROM CLONING TO A COMMERCIAL REALIZATION HUMAN ALPHA INTERFERON

BARON E; NARULA S

SCHERING-PLOUGH RES., 1011 MORRIS AVE., UNION, N.J. 07003.

CRIT REV BIOTECHNOL 10 (3), 1990, 175-190. CODEN: CRDTE

Full Journal Title: Critical Reviews in Biotechnology

Language: ENGLISH

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8/7/12

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7314025 BIOSIS Number: 35094546

MAPPING OF VIRAL EPITOPES WITH PROKARYOTIC EXPRESSION PRODUCTS

LENSTRA J A; KUSTERS J G; VAN DER ZEIJST D A M

INST. INFECTIOUS DISEASES IMMUNOLOGY, FAC. VET. MED., RIJKSUNIVERSITEIT
UTRECHT, P.O. BOX 80105, NL-3500 TD UTRECHT, THE NETHERLANDS.

ARCH VIROL 110 (1-2), 1990, 1-24. CODEN: ARVID

Full Journal Title: Archives of Virology

Language: ENGLISH

8/7/13

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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6817563 BIOSIS Number: 37011942

PROMOTER SPECIFICITY AND MODULATION OF RNA POLYMERASE II TRANSCRIPTION

SALTZMAN A G; WEINMANN R

WISTAR INST., PHILADELPHIA, PA. 19104, USA.

FASEB (FED AM SOC EXP BIOL) J 3 (8), 1989, 1723-1733. CODEN: FAJDE

Full Journal Title: FASEB (Federation of American Societies for
Experimental Biology) Journal

Language: ENGLISH

8/7/14

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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3593556 BIOSIS Number: 23060931

THE SILKWORM BOMBYX-MORI A MODEL FOR MOLECULAR AND CELLULAR BIOLOGISTS

GAREL J-P

DEP. BIOL. GEN. APPL., UNIV. CLAUDE BERNARD LYON-1, F-69622 VILLEURBANNE
CEDEX, FR.

TRENDS BIOCHEM SCI 7 (3), 1982, 105-108. CODEN: TBSCD

Full Journal Title: Trends in Biochemical Sciences

Language: ENGLISH

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8/7/15

DIALOG(R)File 5:BIOSIS PREVIEWS(R)

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3272594 BIOSIS Number: 21064997

BACTERIO PHAGE T-3 AND BACTERIO PHAGE T-7 VIRUS HOST CELL INTERACTIONS

KRUEGER D H; SCHRÖEDER C

INST. VIROL., HUMBOLDT UNIV., DDR-1040 BERLIN.

MICROBIOL REV 45 (1), 1981, 9-51. CODEN: MBRED

Full Journal Title: Microbiological Reviews

Language: ENGLISH

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